REMARKS

Claims 1-27 and 31 are pending in the application. Claims 20-27 and 31 have been cancelled by this amendment. Therefore, claims 1-19 are at issue.

The courteous interview granted by Examiner Bernshteyn to applicants' undersigned attorney and Jürgen Schröder of BASF on June 13, 2007 is hereby noted with appreciation. During the interview, it was agreed that claim 1 would be amended to recite that the superabsorbent particles contain a minimum of about 15%, by weight, clay. The examiner noted that this amendment would distinguish the claims over cited Sun et al. U.S. Patent No. 6,124,391, which discloses a maximum of 10 wt% of an inorganic powder.

During the interview, applicants also thoroughly explained the claimed feature of the clay being "present in the vicinity of the surfaces of the superabsorbent particles." The examiner noted the substantial difference between the structure of the presently claimed superabsorbent particles and the particles disclosed in cited WO 96/30442.

Finally, during the interview, applicants agreed to cancel non-elected claims 20-27 solely to facilitate prosecution. Applicants still consider the restriction requirement as improper. In particular, pending claims 1-27 and 31 have a common technical feature of adding 15% by weight of a clay, or more, to SAP particles during a surface crosslinking process, which is novel and nonobvious. Accordingly, for the reasons set forth in the responses mailed November 22, 2005, March 14, 2006, and September 1, 2006, applicants still contend, per PCT rules, that unity is not lacking between claims of different categories having a common technical feature that is novel and nonobvious.

As stated above, in this amendment, claim 1 has been amended to recite that the surface-crosslinked superabsorbent particles contain a minimum of about 15%, by weight, of a clay. Support for this amendment can be found in original claim 2 and in claim 31.

Claims 20-27 also have been cancelled to delete non-elected claims and facilitate prosecution. Applicants reserve the right to file a continuing application directed to the subject matter of these claims.

Claims 1, 2, 7-10, and 16 stand rejected under 35 U.S.C. §102(b) as being anticipated by WO 96/30442 (WO '442). The basis of this rejection is that WO '442 discloses a water-absorbent polymeric composition containing substantially unreactive water-insoluble particles, e.g., clay particles. For the reasons set forth below, it is submitted that the present claims are neither anticipated by nor obvious over WO '442.

With respect to a rejection under 35 U.S.C. §102(b), applicants wish to point out the following excerpts from MPEP §2131:

"TO ANTICIPATE A CLAIM, THE REFERENCE MUST TEACH EVERY ELEMENT OF THE CLAIM

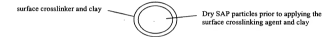
'A claim is anticipated only if each and every element as set forth in the claim is found, either expressly or inherently described, in a single prior art reference.' Verdegaal Bros. v. Union Oil Co. of California, 814 F.2d 628, 631, 2 USPQ2d 1051, 1053 (Fed. Cir. 1987)...'The identical invention must be shown in as complete detail as is contained in the ... claim.' Richardson v. Suzuki Motor Co., 868 F.2d 1226, 1236, 9 USPQ2d 1913, 1920 (Fed. Cir. 1989). The elements must be arranged as required by the claim, but this is not an ipsissimis verbis test, i.e., identity of terminology is not required. In re Bond, 910 F.2d 831, 15 USPQ2d 1566 (Fed. Cir. 1990)."

For the reasons set forth below, it is submitted that WO '442 fails to anticipate the present claims for failing to teach every element of the claims. Additionally, for the reasons set forth below, the present claims would not have been obvious over the cited reference under 35 U.S.C. \$103.

The present claims recite surface-crosslinked superabsorbent particles containing about 15% to about 35%, by weight, of a clay in the vicinity of the surfaces of the superabsorbent particles. These feature are clearly and specifically set forth in claim 1 subparagraphs (ii), (iii)(d), and (iii)(e)., i.e., 1. (ii) about 15% to about 35%, by weight, of a clay, said clay present in the vicinity of surfaces of the superabsorbent particles; 1. (iii) (d) applying a mixture comprising a surface crosslinking agent and a clay to the surface of the superabsorbent particles of step (c) to provide surface-treated superabsorbent polymer particles; and 1. (iii) (e) then heating the surface-treated superabsorbent polymer particles for a sufficient time at a sufficient temperature to surface crosslink the surface-treated

superabsorbent polymer particles and position the clay in the vicinity of the surfaces of the surface-crosslinked superabsorbent particles.

As such, the presently claimed superabsorbent particles have a greater degree of crosslinking at the surfaces of the particles as a result of surface crosslinking in addition to internal crosslinking and the clay is positioned at the vicinity of the surfaces of the particles, i.e., are not distributed throughout the volume of a particle. A claimed superabsorbent polymer (SAP) particle can be visually depicted as follows (not to scale):



Therefore, as recited in independent claim, the clay is added to dry SAP particles (claim 1, subparagraphs (iii)(c) and (iii)(d)). Therefore, as illustrated above, the clay is present in the vicinity of the particle surface only. The benefit of the present invention is that clay on the surface of superabsorbent particles enhances the fluid permeability of the particles and reduces the amount of fine-sized SAP particles.

The superabsorbent particles of the present invention are surface crosslinked. As explained at the interview, surface crosslinked polymer particles are internally crosslinked during polymerization to impart water insolubility, and surface crosslinked after polymerization and drying to improve absorption properties. Surface crosslinked SAP particles therefore have a higher level of crosslinking at the vicinity of particle surfaces compared to uniformly crosslinked particles, i.e., particles that are only internally crosslinked. The reference "Modern Superabsorbent Polymer Technology" illustrates this feature at page 97, figure 3.9, and forwarded to the examiner with the response of December 18, 2006 as Exhibit A. Thus, the polymer particles according of the present invention are different from uniformly crosslinked particles.

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The particles disclosed in WO '442 are substantially different from the presently claimed particles. In particular, WO '442 discloses the use of clay as filler (page 13, lines 1 to 3) to lower extractable levels and improve gel strength (page 13, lines 4 to 6). To achieve these benefits, the clay must be *homogeneously* distributed throughout the polymer particles (as opposed to being present only in the vicinity of the surfaces of the particles). A particle disclosed in WO '442 can be visually depicted as follows:



Example 4 of WO '442 clearly demonstrates that the clay is homogenously distributed throughout the particles. In Example 4, a non-crosslinked polymer (IsobamTM) is mixed with an internal crosslinker (propylene carbonate) and a clay (Bentonite SD-2). The clay is used as a filler and is *homogeneously* distributed throughout the crosslinked IsobamTM polymer particles.

WO '442 discloses IsobamTM and cycloalkylene glycols as suitable copolymers and crosslinkers, respectively (page 7, line 21 to page 8, line 11), and also discloses the *need* of an *internal* crosslinking reaction (page 8, lines 12-14). The three compounds, including the clay, must be *homogenously* mixed before crosslinking. Therefore, the clay in WO '442 *cannot be* present solely at the vicinity of the surfaces of the particles, but rather is distributed throughout the particle, as illustrated above.

Additionally, WO '442 fails to teach or suggest any surface crosslinking.

Accordingly, WO '442 merely discloses uniformly crosslinked particles.

Because WO '442 fails to teach each and every feature recited in the claims, WO '442 cannot anticipate the present claims (MPEP §2131). In addition, the differences between the WO '442 disclosure and the present claims are substantial and nonobvious differences.

As illustrated above, WO '442 discloses particles having a clay distributed throughout the particles to improve gel strength and reduce extractables. WO '442 also fails to teach or suggest any surface crosslinking. In contrast, the present claims recite surface crosslinked particles having a clay present in the vicinity of the particle surfaces. WO '442 fails to provide any motivation or apparent reason for a person skilled in the art to modify the WO '442 teachings in the radical manner required to arrive at the presently claimed invention. Furthermore, contrary to the examiner's contentions, WO '442 does not teach a "substantially identical" polymer to the claimed particles. WO '442 does not teach a surface crosslinking agent, and WO '442 teaches a substantially different process for producing substantially different superabsorbent particles, i.e., teaches a homogenous incorporation of the clay in the particle.

Accordingly, it is submitted that claims 1, 2, 7-10, and 16 are neither anticipated by, nor obvious over, WO '442, and that the rejection should be withdrawn.

Claims 1-5, 7-17, and 31 stand rejected under 35 U.S.C. §103 as being obvious over Sun et al. U.S. Patent No. 6,124,391 ('391) in view of WO '442. Claim 6 stands rejected under 35 U.S.C. §103 as being obvious over the '391 patent in view of WO '442. Claims 18 and 19 stand rejected under 35 U.S.C. §103 as being obvious over the '391 patent and WO '442 in view of Beerse et al. U.S. Patent Publication 2002/0006886 ('886). In view of the amendments to the claims, for the reasons set forth above, and for the reasons set forth below, it is submitted that these rejections are in error and should be withdrawn.

The '391 patent discloses the preparation of dried SAP particles, admixing the dried SAP particles with 0.2 to 10 wt% of an inorganic powder and a surface crosslinking agent (column 7, lines 27-32 and 43-45), and heating the resulting mixture. The inorganic powders can be a clay. The inorganic powder can be added to the SAP particles before, during, or after the surface crosslinking step. Importantly, the '391 patent specifically discloses that the clay is added to the SAP particles in an amount sufficient to achieve anticaking properties, up to a maximum of 10 wt%, and preferably less than 10 wt%, (see '391 patent, column 7, lines 27-32). Additionally, the examples of the '391 patent are SAP particles having a clay concentration of 0.5 to 3 wt%.

As stated above, the '391 patent is directed to incorporating an anticaking and dedusting amount of an inorganic powder to SAP particles. The *maximum* amount of inorganic powder added to the SAP particles, as disclosed in the '391 patent, is 10 wt%. In

contrast the present claims specifically recite a *minimum* amount clay of about 15%, by weight.

In addition, and as set forth in the Amendment of September 1, 2006, the differences between the '391 patent disclosure and present claims are nonobvious differences. The '391 patent discloses amounts of inorganic powder typically used in the art to impart anticaking and dedusting properties to SAP particles. The '391 patent contains no teaching or suggestion that would motivate a person skilled in the art to increase the amount of inorganic powder above the disclosed maximum limit of about 10%, by weight. In fact, persons skilled in the art would have had no incentive to increase the amount of inorganic powder about 10 wt%, because the '391 patent teaches that dedusting is achieved at inorganic powder amounts well below 10 wt% (e.g., see '391 examples). Therefore, persons skilled in the art would consider using any amounts of clay above 10 wt% as being wasted.

However, applicants have found that including a clay in the surface crosslinking step, in the claimed amount of about 15% to about 35%, by weight, provides the fluid unexpected benefits of reducing the amount of fine-sized SAP particles and improving the fluid permeability of the clay-treated SAP particles. See specification, page 7, lines 32-35, and page 8, lines 12-16, for example. The reduction of fine-sized SAP particles is a long felt need in the art because fine-sized particles have invariably been generated during SAP production, and either are recycled in a time consuming, expensive process step or are discarded as waste.

Applicants further have provided objective evidence demonstrating the new and unexpected results provided by the presently claimed invention. In each of Examples 5 and 6, at pages 30-33 of the specification, applicants provide a series of SAPs containing varying amounts of clay added to the SAP during the surface crosslinking step. Example 5, in the table at page 30, shows that an amount of clay disclosed in the '391 patent, i.e., 5% and 10 wt%, generates 20% and 16 wt%, respectively, of SAP particles having a diameter less than 200 μ m (i.e., fine-sized SAP particles). By increasing the amount of clay to 15 wt% in accordance with presently claimed invention, fine-sized particles having a diameter less than 200 μ m drops substantially to 4.5 wt%. Further increasing the amount of clay up to 35 wt% results in further decreases in the amount of fine-sized SAP particles. Applicants claims are now tailored to the unexpected results provided in the disclosure (see Examples 5 and 6), which clearly distinguishes the present claims over the '391 patent.

The table of Example 5 also shows a substantial increase in fluid permeability in SAP particles having 15 wt% or more clay (inventive) over SAP particles having 5 wt% or 10 wt% clay (comparative and within the '391 disclosure). See SFC data in the table of Example 5. The importance of a high SFC value as relating to SAP particle permeability is discussed at page 28, line 17-26 of the specification. The table of Example 6 of the specification contains similar results with respect to both reducing the amount of fine-sized SAP particles and improving SAP particle permeability when a clay is present in an amount recited in the claims. Notably, the SAP particles also performed well with respect to absorption properties even in the presence of high amounts of clay (see specification, page 33, lines 1-3 and AUL and CRC data in the tables of Examples 5 and 6).

Contrary to the examiner's statements in the Office Action, these results are unexpected in view of the teachings of the '391 patent, which are limited to a maximum of 10 wt% inorganic powder added to SAP particles to provide anticaking and dedusting properties. The '391 patent absolutely fails to teach or suggest using greater amounts of an inorganic powder for any purpose. The '391 patent teaches that the amount of clay to be used is typically about 0.2% to about 10%, more preferably about 0.5% to about 7%, and even more preferably about 0.9% to about 5.5%. The teaching of the '391 patent is that the best results can be expected in the most preferred range from 0.9% to 5.5%. Thus, one of ordinary skill in the art would not have been motivated to use clay in an amount substantially greater than that in the '391 patent disclosure. In fact, the '391 patent discourages increasing the amount of inorganic powder added to the SAP.

Persons skilled in the art certainly would have had no incentive from the '391 patent (directed to anticaking and dedusting) to increase the amount of clay added to SAP particles with any reasonable expectation of substantially reducing the amount of fine-sized SAP particles and improving SAP particle permeability, while also retaining SAP particle absorption properties.

WO '442 does not overcome the deficiencies of the '391 patent. First, WO '442 discloses particles substantially different from the presently claimed particles, as discussed above, i.e., clay distributed throughout the particle and no surface crosslinking. Second, after reading the '391 patent, a person skilled in the art would not have been motivated to increase the amount of clay to greater than 10%, as discussed above. Although the type of particle disclosed in WO '442 may be able to incorporate up to 25% clay (because

the clay is distributed throughout the particle), it does not correlate to the amount of clay can be present in the type of particle disclosed in the '391 patent. In fact, the '391 patent discourages such an increase in the amount of clay at the surfaces of the particles.

Therefore, for the reasons set forth above, it is submitted that claims 1-17 would not have been obvious over the '391 patent in view of WO '442 under 35 U.S.C. §103.

With respect to claims 18 and 19, these are additional embodiments of the present invention and are patentable for the same reasons presented above for claims 1-17. In particular, applicants do not rely solely upon the features recited in claims 18 and 19 for patentability, but rely on *all* the features recited in these claims, including the features of claim 1 and 15 from which claims 18 and 19 depend.

Furthermore, applicants maintain that the '886 publication does not overcome the deficiencies of the '391 patent and WO '442. The '886 publication merely discloses well known quaternary ammonium compounds (QACs). The QACs of the '886 publication are *not* incorporated into clay particles, but are added *directly* to "a water insoluble substrate comprising a nonwoven layer" (see paragraphs [0010], [0023] through [0051], and [0247]). A clay is *not* a nonwoven layer as disclosed in the '886 publication. The '886 patent fails to teach or suggest any combination of a QAC and a clay to provide an organophilic clay as required in claims 18 and 19, and furthermore the '886 publication is not remotely directed to SAP particles.

In fact, the QACs of the '886 publication are disclosed as optional, nonlathering cationic surfactants. The disclosure relating to QACs is nothing more than a general description of available QACs. At most, the '886 publication discloses that both an SAP and a QAC can be optional ingredients in a composition disclosed therein. There is no teaching that an SAP and a QAC should both be present, and no teaching or suggestion of including a clay. Furthermore, persons skilled in the art would not add a hydrophobic QAC to an SAP because the absorbency properties of the hydrophilic SAP would be adversely affected.

Therefore, persons skilled in the art would have had absolutely no incentive to consider combining any teaching from the nonanalogous '886 publication with the '391 patent and WO '442 with any reasonable expectation of providing a useful superabsorbent polymeric particles. Further, even if such a combination of references was considered, the combination would not lead a person skilled in the art to incorporate a standard QAC of the '886

publication (taught for use as a surfactant) into a clay (which ties up the QAC), then incorporate such a treated clay in the presently claimed amount into SAP particles during a surface-crosslinking step. Such jumps in reasoning to arrive at the presently claimed invention are neither apparent nor remotely suggested by the combination of the '391 patent, WO '442, and the '886 publication.

To support the rejection, the examiner relies upon a definition of the term "tallow." It is well known that a QAC can have a tallow component, but the '886 publication is limited to disclosing a QAC, not even remotely addressing an organophilic clay having a QAC component. An organophilic clay has properties different from an untreated clay because of the QAC. The QAC in the organophilic clay performs differently from a free QAC as disclosed in the '886 publication. The QAC in the '886 patent is used to provide a nonlathering surfactant. In the present claims, the QAC is *not* available to act as a surfactant, but modifies the properties of the clay by substituting organoamminium ions for metal cations present between the clay layers (see specification, page 23, lines 23-29). Applicants do not positively recite that the QAC is "bound to a clay" because persons skilled in the art are well aware of the structure of an organophilic clay. Also note claim 15 and the specification at pages 24 and 25, so no new matter issues exist as suggested by the examiner.

The examiner further misunderstands the claimed invention vis-à-vis the '886 publication by statements in the Office Action. The examiner's conclusory statement that it would have been obvious to "incorporate organophilic clay selected from tallow" into an SAP to obtain additional functional group linkages is totally incorrect. These hydroxyl and amino functionalities may be present in a composition of the '886 publication because the QAC is free, and is not bound to a clay. In an organophilic clay, the QAC is bound and is not available to provide any such functionalities. Further, if such functionalities were available, they may be detrimental and adversely affect the absorbency of the SAP particles.

Applicants reiterate that the '886 publication is nonanalogous art. The examiner previously admitted that the '886 publication is directed to a dry article for personal cleansing, and more generally to an article that *delivers* a compound having a benefit. An SAP does *not* deliver compounds, but rather absorbs aqueous solutions. The goal of the '886 patent teachings would be frustrated by using a strongly absorbent SAP in place of the nonabsorbing nonwovens disclosed in the '886 patent. The SAP absorbs, whereas the nonwovens deliver.

The examiner's reliance on *In re Spada* also is misplaced. Applicants are not reciting *properties* in claim 1. To the contrary, applicants are reciting *structure*, i.e., clay positioned in the vicinity of the surfaces of a dry superabsorbent polymer particles during surface crosslinking.

The examiner's statement that the references suggest increasing the amount of clay in a particle "to receive economic saving" and "there is little or no deleterious effect on other properties" is inapposite. Taken to a logical conclusion, a clay can be a *total* substitute for superabsorbent polymer particles in order to attain economic savings, but the resulting product would be essentially useless for absorbing liquids compared to superabsorbent polymer particles. Also, although WO '442 discloses no deleterious effect in properties, the reference also shows *no improvement* in properties. The presently claimed superabsorbent polymer particles have a structure different from those in WO '442 and showed improvements as described above when present at about 15%, as claimed and as demonstrated in Examples 5 and 6 discussed above. In addition, in view of the substantial structural differences between the presently claimed particles and WO '442, a direct comparison is meaningless. In fact, applicants have provided data showing the particles having 5% or 10%, by weight, clay at the particle surface are inferior to particles having 15%, by weight, clay at the particle surfaces (see Examples 5 and 6).

For all the reasons set forth above, it is submitted that claims 1-19 would not have been obvious over the '391 patent in combination with WO '442, or in further combination with the '886 publication, and that the present rejections should be withdrawn.

The pending claims are in a form and scope for allowance. An early and favorable action on the merits is respectfully requested.

Should the examiner wish to discuss the foregoing, or any matter of form in an effort to advance this application toward allowance, the examiner is urged to telephone the undersigned at the indicated number.

Dated: July 16, 2007

Respectfully submitted,

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